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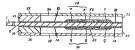
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A biosensor and a method for measuring a concentration of a substrate in a sample (54) (57) The present invention provides a biosensor

comprising en electrical insulating substrate having a plurality of surfaces, a plurality of electrode systems formed on at least two of the plurality of surfaces of the substrate, each of the electrode systems having a working electrode and a counter electrode, and a plurality of reaction layers provided on at least two of the plurality of surfaces of the substrate, each of the reaction lavers Fig. 12

containing an exidereductase, wherein the plurality of electrode systems are respectively provided on different surfaces of the substrate, the reaction layers are provided on the surface on which the plurality of electrode systems are formed, and the kind of oxidoreductases contained in the plurality of reaction layers are different from each other



Description

BACKGROUND OF THE INVENTION

1 Field of the Invention:

The present invention relates to a biosensor according to claim 1 and a method for measuring the concentrations of at least two substrates (specific components) in a sample by using the biosensor.

2. Description to the Prior Art:

Various types of blosensors utilizing specific catalyses of enzyme have been recently developed. A socharide biosensor will be described as an example of such biosensors as follows:

The optical rolation method, the colorimetic method, the roducinety method and other methods using different fixths of drometographies have been as developed are methods for quantitative analysis of searchardes, However, none of these methods can provide high accuracy due to the releasely live specificity against sectharides. Additionally, the optical rolation method is easy to penaltho bit is supply influenced by a the operating temperature. Therefore it is not appropriate to be common used an thome and the list.

The sexchandes contained in faul are generally sesseed as succhandes contained in faul are generally sesseed as succhande degrees. Are fercionered of the sight reflaction system is other used for quantifying the 30 socialization system in ordinary services by using the system of the second system of the syste

A glucose sensor will now be described as an 40 example of a biosensor used in a clinical field.

A conventional method for quantifying glucose con-

tained in blood is to centrifuge blood taken from a patient and then to measure the thus obtained blood plasma. This method requires a lot of time as well as labor. Therefore, a censor that can directly measura glucose in blood obtained from a patient is desired.

A sonor similar to a teel paper for unanalysis has been developed as similed gloose servic. This glucose service comprises a support in a stick shape and a hotel fixed in the support. The hotel reliables is an hotel fixed in the support. The hotel reliables is supported by the support of the support of energies resident. Belood is dropped on the support of the glucose service and the chemps of the color of the glucose service and the chemps of the color of the dropped by desired and support of the disporting to steasily desired or optically reassured, whereby the control of the support of the support of the support of the support of the disporting to steasily desired or optically reassured, whereby the cover, the quartifiers method using the success service were the quartifiers method using the success service. has low accuracy due to interference by the colored

materials in the blood. EP 0 359 831 A1

discloses the following glucose sensor with high eccuracy se a method for quantifying a specific component in a sample liquid from a living body such as blood without diluting or string the sample liquid:

The biosensor comprises a base 42, and a spacer 3 and a cover 4 that are lamineted integrally onto the

3 and e cover 4 that are lamineted integrally onto the base 42 es is shown in Figures 13 and 14.

The base 42 comprises an electrical insulating selsstrate 1, an electrode system 43 formed on the substrate 1 by screen printing, etc. and a reaction layer 44 provided on the electrode system 43. The electrode system 43 includes a working electrode 45 and a counter electrode 46 that are electrically insulated from each other by an insulating layer 47. The working electrode 45 and the counter electrode 46 are connected to leads 12 and 15 formed on the substrate 1, respectively.

The reaction layer 44 includes a hydrophilic polymer, an axidoreductase and electron ecceptors and covers the working electrode 45 and the counter electrode 46.

trode 46. As is shown in Figure 13, the spacer 3 is in a Ushape and has a groove 17. When the spacer 3 and the cover 4 are laminated on the base 42, e peasage 18 through which a sample liquid passes is formed between the base 42 and the cover 4 es is shown in Figure 14. One end of the passage 18 is open at one end of the base 42 and the coveres as a sample sup-

by but also was me opening serves as a sample supply port 23. The other end of the passage 18 is open the cover 4 and the opening serves as an air port 24. The operation of the glucose service with the abovementioned structure is an follows: A sample liquid supmentioned structure is an follows: A sample liquid sup-

s pied through the sumple supply port 22 neaches the reaction layer 44 through the passage 18 and the oxidoreductase and the electron acceptors contained paid. Thus, while an enzyme reaction is proceeded between a substrate in the sample liquid and the coldoreduction, the electron acceptors are reduced. After trishing the enzyme reaction, the reduced electron acceptors are

electrochemically oxidized. A value of an oxidation current obtained at this point provides a concentration of at the substrate in the sample liquid. However, the conventional biosensor has the following disadvantages:

The sample liquid may include reductive materials that can reduce the electron acceptors other than the substante to be measured. Moreover, viscoeity, etc. of the sample liquid to be measured very.

Accordingly, in measuring a concentration or the substrate in the sample figuid including the substrate and another roductive material that can reduce the elecstron acceptors, the response values of the sensor are inconstant, and therefore, the reductive material should be eliminated before the measurement. Such a pretreament results in increasing the number or steps in measuring the concentration of a substrate in a sample liquid.

Moreover, the sensor response elso depends upon a measuring time. For example, an accurate concentration can not be obtained when the oxidation current is measured before completely finishing the reaction.

Furthermore, time required for the sample liquid to reach the reaction layer and the rate of the reaction of the substrate to the enzyme depends upon the viscosity of the sample liquid. Therefore, the inconstant viscosities result in inconstant sensor response.

SUMMARY OF THE INVENTION

The biosensor of this invention comprises an electronia makening, substante having a pursible of verticese, is plurally of electricise systems for medio on a least two of a plurally of electricise systems for medio on a least two of the electricise systems for medio on the electricise systems of electricise systems for the plurally or attacked and as an occurried reflection, and a plurally of reaction layers provided on all leasts two or the plurally or attacked on the substants, each of the reaction layers containing an octation of the electricise of the substants of the subterior of the electricise of the substants of the electricise of the substants of the electricise of the subtraction. In the electricise of the subtraction of the local of devolved-classes contained in formed, and the local of oxidoved-classes contained in formed, and of the local of oxidoved-classes contained in formed, and of the local of oxidoved-classes contained in the measurement of the local oxidoved-classes contained in the measurement of the local oxidoved-classes contained in the measurement of the local oxidoved-classes contained in the local oxidoved-classes contained in the local oxidoved-classes contained in the local oxidoved-classes contained in

Since the kinds of axidoreductases contained in the reaction layers provided on respectively different surfaces of the substrate are different from each other, it is possible to prevent various components such as butter components, enzymes and the life in the respective reaction layers from moving towards each other.

As a result, when a sample liquid is supplied to the sensor of the invention, the concentration of substrates contained in the sample liquid can be measured in each reaction law. Thus, the precision in measuring the note

centration of the substrate can be improved.

These and other advantages of the present invention will become apparent to those skilled in the art upon reading and understanding the following detailed description with reference to the accompanying figures.

Figure 12 shows e preferred embodiment of the present invention, whereas on figures 1 to 11 additional aspects of the invention are displaced.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a plen view of e base of a glucose sensor eccording to an example the present invention con-

Figure 2 is an exploded perspective view of the glucose sensor of Figure 1 from which a reaction layer is removed.

Figure 3 is a sectional view of the glucose sensor of Figure 1.

Figure 4 is a plan view of a base of a glucose sen-

sor according to another example the present invention concerns.

Figure 5 is a plan view of a base of a glucose sensor according to still another example the present invention concerns.

Figure 6 is an exploded perspective view seen from one side of a glucose sensor according to still another example the present invention concerns.

Figure 7 is an exploded perspective view of the glucose sensor of Figure 6 seen from the other side. Figure 8 is a plan view of a base of the glucose sen-

sor of Figure 6.
Figure 9 is a sectional view of the glucose sensor of as Figure 6.

Figure 10 is a plan view of a base of a glucose sensor according to still another example the present invention concerns. Figure 11 is a sectional view of a glucose sensor

30 according to still another example the present invention concerns.
Figure 12 is a sectional view of a saccharide sensor

according to an example of the present invention.

Figure 13 is an exploded perspective view of a conventional disposable glucose sensor from which a reaction layer is removed.

Figure 14 is a sectional view of the glucose sensor of Figure 13.

DESCRIPTION OF THE PREFERRED EMBODI-MENTS

Throughout the drawings mentioned in the following description of the examples, the same element has a common reference numeral. Per of the description is omitted as occasion demands,

Example 1

A glucose sensor will now be described.

The glucose sensor comprises a base 2, and a spacer 3 and a cover 4 integrally laminated on the base 2 as is shown in Figures 2 and 3.

The base 2 comprises en electrical insulating subsimal 1 made from polyethylene terephthalate, and trode system formed on the subsette 1 by ecreen printing and the like. The electrode system comprises a main electrode system 19 and a sub-electrode system 20 provided on the substrate 1 with an interval therebetween. The main electrode system 19 and the sub electrode system 20 include working electrodes 6 and 8 and counter electrodes 7 and 9, respectively. The working electrodes 6 and 8 and the counter electrodes 7 and 9 are electrically insulated from each other by an insulatificial lower 10.

A lead 12 formed on the substrate 1 is electrically connected to the working electrode 6 Of the main electrode system 19, e lead 13 to the counter electrode 7 of the main electrode system 19, e lead 14 to the working electrode 9 of the sub-electrode system 20, and e lead 15 to the counter electrode 9 of the sub-electrode system 20.

A reaction layer 5 covers the working electrode 6 for the counter electrode 7 of the main electrode system 19. The reaction layer 5 includes carboxy methyl cellulose (hereinatter called the CMC) as a hydrophilic polymer, glucose oxidas (ECL1.3.4, the releaster called the GOD) as an oxidoreductase, and potassium terricy-

anide as dedoom ecceptors. As is shown for plane 2 in second in a billing of the plane 2 in second 2 in second 2 in a billing of the plane 2 in second 2 in a plane 2 in a manded on the base 2, a passage 18 in through which a cample legical passess is somed between the base 2 and the cover 4. One of the passage 18 is good or one supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a plane 2 in a plane 2 in a supply plane 2 in a s

The glucose sensor was produced as fallows: Silver paste was printed on the authorize 1 by means of screen printing to form the leads 12, 13, 14 and 18, 35. Then conductive cushop pasts including reals binder was printed on the substrate 1 to form the working electrodes 6 of the main electrode system 19 and the working electrodes 6 and the counter electrode 9 of the sub-silver binder pasted on the substrate 1 to form the working electrode 5 and the counter electrode 9 of the sub-silver binder system 30.

The working electrodes 6 and 8 and the counter electrode 9 were electrically connected to the leads 12, 14 and 15, respectively.

Next, insulating purse was printed on the substants of to form the insulating layer 10. The insulating layer 10 and occurred report of the concerned the peripheral portion of the working electrode 6 was exposed. Further, the insulating layer 10 covered 6 was exposed. Further, the insulating layer 10 covered or exert of the factor 12, 13, 14 and 15, respectively. The working electrode 9 and the countries electrode 9 and the continuation layer 10 covered on the continuation layer 10 on the 19 prefettering area of the continuation layer 10 on the 19 prefettering area of the working electrode 5 and the counter electrode 9 were respectively exposed.

Then, conductive carbon peate including resin as binder was printed on the insuleting layer 10 so as to come in contact with the load 13 to form the counter electrode 7 of the main electrode system 19. The base 2 shown in Floure 1 was produced in this manner. Next, an aqueous solution including the GOD as the outdoneductuse, potassiss metriograndie as the electron ecceptors and the 0.5 wt% of CMC as the hydrophilic polymer was dropped on the working electrode 8 and the counter electrode 7 of the main electrode system 19, and was chief of the warmer's chief at temperature of 50°C for 10 minutes to form the reaction layer 6. The reaction layer 6 and the easily formed to easily formed in the grant of the system 19.

Then, a mixed aqueous solution including potassium ferricyanide and the CMC was dropped onto the working electrode 8 and the counter electrode 9 of the sub-electrode system 20 and dried to form e reference layer 25.

After forming the reaction layer 5 and the reference layer 25 on the substants 1, the cover 4 and this spacer 3 were lamineted to be athered onto the base 2 as shown in Figure 2 with disshed lines. Thus, the passage 18 having a comparatively amel cross section was formed which was defined by the groove 17 of the spacer 3, the cover 4 and the base 2.

The thus produced glucose service was supplied with 3 yet of mixed queues earlier inclusing glucose and seconds caid see a sample lagoid through the service and seconds caid see a sample lagoid through the service lagoid, or the service lagoid, or the service lagoid, or the service lagoid or the service lagoid covering in content with the service supply or 23 after big or the sector is involved in the pear-point 28 after big or the sector is involved in the pear-point 28 after big or the sector is involved in the pear-point 28 after big or the sector is involved in the pear-point 28 after big or the sector is involved and the reached that the sector is the sector is the pear-point 28 after big or the sector is required to the pear big or the sector is the sector of the sector is the sector of the sector

content with the sample supply profit of The sample size plant plant of the profit of the same described system of the size possible, see that the same described system of the size plant size showing the size of the size plant size were respectively dissolved in the sample legal of size of the size of the size of the size of size of the size of the size of the size of si

The potassium ferriogranide on the sub-electrod system 20 was reduced by ascorbic ecid in the sample system 20 was reduced by ascorbic ecid in the suplequid to generate potassium ferrocyunide. The oxidation current value [a obtained by the application the above-mentioned productermined voltage was caused by oxidation of the potassium intercopyratide. Therefore, the oxidation ourrent value i_{ij} was in proportion to the amount of ascorbic edid containing in the sample light.

A voltage of +0.5 V on the basis of the voltage at the counter electrode 7 of the main electrode system 19 was applied to the working electrode 6 of the main elec-

was measured.

trode system 19 one minute after detecting the above described impedance change. An exidation current value I₁ of five seconds after the application was measured.

The exidation current value I_t is a total value of e sourrent caused by exidation of potassium ferrocyanide generated by reduction by escorbic exid and one caused by exidation of potassium ferrocyanide generated in exidating plusoes by the GOD.

When a coefficient for correcting the difference in the responses in both the electrode systems is taken as k, a current value represented as I₁-40, dotasty corresponded to the glucose concentration in the sample liquid

Next, responses obtained in the following operations (1) and (2) were compared, using thirty glucose sensors.

(f) After supplying the sample liquid to the sensor through the sample supply not, a sufficient supply of the sample liquid to the reaction layer 5 was delected by delecting the change of the electrical channel residence of the electrical system. Then the sensor responses were obtained in the above-mentioned resemble.

(2) Instead of the detection of the change in the electrical characteristics, sufficient supply of the sample liquid to the reaction layer 5 was confirmed as by visual observation. Then the sensor responses were obtained by measuring the addition currents was to find in the same manner as above.

As a result, e coefficient of variation (a CV value) as showing the dispersion of the responses was 2% in operation (1) and 5% in operation (2).

As meritioned above, the coefficient of the variation in operation (1) was realizer than that in operation (2). This scene to be because the inconstancy in time from 40 the measurement of the response current in the subdistincts system to the list the main electrodic system was made amaller in operation (1).

In this example, the GOD in his reaction layer is used intermobilities in the main electricide system 19, However, times the reaction layer for contained the hydrophilic way. Except the sample liquid was proposed to the control liquid was sometimed to the containing in the carried layer for supervised on their the sample liquid. Therefore, depending on instantial the materials did not move broad the sub-decidated in the carried layer for supervised on their the materials did not move formed the sub-decidated spectra 20 in a level product of time. A more accounter measurement can be siftedly statistical by immobilizing an entire that the GOD in the man electricid get entire than the GOD in the man electricide get an entire was such as the GOD in the man electricide.

Example 2

Figure 4 is a plan view of a base 2 of a glucose sensor produced as another example of the biosensor the present invention concerns. This glucose sensor was produced as follows:

Silver paste was printed on en electrical insulating substrate 1 made from polyethylene terephthaliate by screen printing to form leads 12, 13, 14 and 15. Next, conductive carbon paste including resin binder was printed on the substrate 1 to form a working electricide of e main electricide system 19 and a working electricide.

8 of a sub-electrode system 20.

The working electrodes 6 and 8 were electrically connected to the leads 12 and 14 reconcises.

combination to the leader 12 and 14, respectively. Insulating paster was then printed on the substrate 1 to form an insulating layer 10. The insulating layer 10 covered perspheral portions of the working electrodes 6 end 8 so that a predetermined area of the working electrodes 6 and 8 was exposed, respectively. Moreover, the insulating layer 10 covered or part of the leads 12, 13, 14 and 15, respectively.

Next, conductive carbon paste including resin binder was printed on the insulating layer 10 so as come in contact with the leads 13 and 15 to form a counter electrode 7 of the main electrode system 13 and e counter electrode 9 of the sub electrode system 20. In this manner, the base 2 shown in Figure 4 was

produced. A related equations solution including the GOO, A related equations solution including the GOO, A related equation to the solution of the solution o

respectively.

Then a spacer 3 end a cover 4 were laminated integrally on the base 2 having the reaction layer and the reference layer in the same manner as in Example 1, to form the ducose sensor.

Since the reference layer comprising the BSA, potassium ferricyanide and the CMC is formed on the sub-electrode system 20, conditions for diffusion of a preductive material such as escorbic ecid on the sub-electrode system 20 and the like, are similar to those on

the main electrode system 19.

In the case that proteins such as the GOD and the BSA exist on the electrode systems, the activity of the electrode may be partly degraded by absorption of such proteins. However, a layer including proteins proteins proteins believed by the electrode system of such proteins. However, a layer including proteins provided on both the main and such electrode systems or the such proteins and such proteins are not such as the such electrode systems of the such electrode systems.

tern due to the absorption of the proteins. As a result. the correction between the oxidetion current values in

both the electrode systems 19 and 20 can be simplified. Such an advantage is especially remarkable in a sensor ueing carbon es e main electrode material.

Example 3

Next, a fructose sensor as an exemple of e biosensor will be described.

Figure 5 is a plan view of e base 2 of a fructose sensor produced as still another example of the biosensor eccording to the present invention. The fructose sensor was produced as follows:

As it shown in Figure 5, silver paste was printed on an electrical insulating substrate 1 made from polyethylene terephthalate by screen printing to form leads 12, 13 and 14. Next, conductive carbon paste including resin binder was printed on the substrate 1 to form a working electrode 6 of a main electrode system 19 end a working electrode 8 of e sub electrode system 20. An insulating layer 10 was then formed on the substrate 1 by using insulating paste. The insulating laver 10 covered propheral portions of the working electrodes 6 and 8 so that a predetermined area of the working electrades 6 and 8 was respectively exposed. Further the insulating layer 10 covered a part of the leade 12, 13 and 14, respectively.

Next, conductive carbon paste including resin binder was printed on the insulating layer 10 so as to come in contact with the lead 13 to form a counter electrode 7. Thus the base 2 was produced.

Next, a mixed aqueous solution of fructose dehydrogenase (EC1,1.99.11: hereinafter called the FDH) es an oxidoreductase, potassium ferricyanide es electron acceptors and the CMC as a hydrophilic polymer in e phosphate buffer solution (pH = 5) was dropped on the working electrode 6 and the counter electrode 7 of the main electrode system 19, and dried in a warm-air drier at e temperature of 40°C for 10 minutes to form a reaction layer (not shown).

A spacer and a cover were laminated integrally on the thus obtained base 2 in the same manner as in Example 1 to produce the fructose eensor.

Three ut of a mixed aqueous solution of fructose 46 and escorbic acid was supplied to the fructose sensor. A voltage of +1 V on the basis of the counter electrode 7 was applied to the working electrode 8 of the sub electrode system 20 end an codation current value le was measured. Since no axidoreductase end electron ecceptors existed on the working electrode 8 of the subelectrode system 20, the current value Is was an coldation current value of ascorbic acid contained in the sample liquid. Mareover, since no hydrophilic polymer end the like for preventing diffusion of materials existed on the sub electrode system, on accurate current value In was obtained immediately after the supply of the sample liquid. The current value In was in proportion to the scorbic acid concentration.

Moreover, a voltage of +0.5 V on the besis of the counter electrode 7 was applied to the working electrode 6 of the main electrode system 19 one minute after the supply of the sample liquid to the sensor through the sample supply port. A current value I, of 5 seconds after the application was measured. The current value I, is a total value of an oxidation current of potassium ferrocyanide generated by e reaction with escorbic ecid and one generated in a reduction of fructose by the FDH.

The amount of ascorbic acid contained in the sam ple liquid was quantified by using the current value In. A fructose concentration in the sample liquid was calculeted from the quantified amount of ascorbic ecid and the amount of potassium ferrocvanide quantified with the current value 1.

in the fructose sensor eccording to this example. the counter electrode 7 serves as a common counter electrode to the main and sub electrode systems 19 and 20. The production of the sensor can be simplified by using the counter electrode in common in this manner. Moreover, the cost for producing the sensor can be reduced by using one less lead. In addition, increasity of the surfaces of the electrode systems on the substrate 1 can be minimized, thereby preventing the reaction layer from peeling off from the electrode systems. As e result, e sensor with excellent conservative and stable properties can be produced, and an accurate sensor response can be obtained by making the movement of the sample liquids smoother on the electrode systems.

Example 4

A method for measuring e glucose concentration in whole blood by using the glucose sensor produced in the same manner es in Example 1 will now be described

Whole blood was supplied to the glucose sensor through the sample supply port 23. The entire glucose sensor including the reaction layer 5 was in a dry condition before the supply of the sample liquid. The sample liquid, that is, the whole blood reached the sub electrode system 20 tirst, thereby reducing an impedance between the working electrode 8 and the counter electrode 9 in the sub electrode system 20. The impedance change was detected through the leads 14 and 15.

Next, the whole blood reached the main electrode system 19. When the reaction layer 5 on the main electrode system 19 was dissolved, an impedance between the working electrode 6 and the counter electrode 7 of the main electrode system 19 was reduced. The impedence change was datected through the leads 12 and 13.

When the reaction layer 5 was dissolved in the whole blood, the glucose in the blood was oxidized by the GOD, and at the same time potassium ferricyanide was reduced into potassium ferrocyanide. One minute after the supply of the whole blood to the glucose sensor, a voltage of +0.5 V on the basis of the counter electrode 7 was applied to the working electrode 6, and an oxidation current 5 seconds after the application was measured. The obtained current value was due to the reduction of potassium ferrocyanide and was in proporfilion to the concentration of glucose, the substrate to be measured.

When the above denorbled oxidation current values were measured by using the whole blood sample with a hemotroir of 20% to 50%, a higher hematocrit reduced the exidation current value. Moreover, when the difference of time required for detecting the impedance of time required for detecting the impedance of time required for detecting the impedance of the produced time to the produced of the produced or time to the produced of the produced or time to the prod

whole blood with a hematocrit of 20% to 60%. When values obtained by correcting the coldation current value with the above factor twen taken as sensor responses, the sensor responses were constant regardless of the hematocrit value in the whole blood sample end corresponded to the glucose concentration

In this example, the sample liquid can be explicit through the in prof. A using the sample septy port 23 as an air port. In this case, the impodance change is first detected in the mis electrode system, and then in the sub-electric system. The difference in time 12, serequired for detecting the impodance change between the main electrode system and the sub-electrode system and the sub-electrode system and the sub-electrode system and the sub-electrode system the main electrode system and the sub-electrode system than does not reconstruct or sub-electrode system and the sub-electrode system than the sub-electrode system to the sub-electrode system that the sample of the sub-electrode system that the sub-electrode system

Example 5

in the whole blood

A glucose sensor will now be described. Figure 8 is an exploded perspective view of the glucose sensor seen from one side from which e readilon layer 50 is removed. Figure 7 is an exploded perspec-

tive view of the glucose sensor seen from the other side from which e reaction layer 50 is removed. Figure 8 is a 40 plan view of a base 2 of the glucose sensor produced in this example. Figure 9 is a sectional view of the glucose sensor produced in this example.

A method for producing the glucose sensor will now be described.

Sever paste was printed on a substrate 1 made.

Sever peate was printed on a substate 1 mode from polythylines in septembhalis by screen printing to from leaks 12 and 13. Med., conductive carbon paste form leaks 12 and 13. Med., conductive carbon paste including risk bindwes perifical on the substate 1 to form a working electrode 6 of a main electrode system 19. The working electrode 6 treat place training 19. The working electrode 6 was electrode as substate 1 to substate 1 to form on insulating layer 10. The insulating substate 1 to form on insulating layer 10. The insulating electrode 6 to that a prodetermined even of the working electrode 6 to that a prodetermined even of the working electrode 6 was exposed.

Next, conductive carbon paste including resin binder was printed on the insulating leyer 10 so as to come in contact with the lead 13 to form a counter electrode 7 of the main electrode 19.

On the reverse surface of the insulating substants I on which the above-mentioned electrode pattern was printed, leads 14 end 15, a sub-electrode system 20 (including a working electrode 8 and a counter electrode 9) end an insulating layer 16 were formed by printing, thereby producing the base 2 as shown in Figures 7 and

The structures of the main electrode system 19 and the sub electrode system 20 were the same, and therefore, the areas of the working electrodes 6 and 8 were also the same.

elso the same.

A mixed aqueous solution including the GCD as the calcioreductate, potassium ferricyanide as the electron acceptors and the CMC as the hydrophilic polymer was dropped on the main electrode system 19, and dried in a warm-six order at a temperature of 5°CC for 10 minutes.

Next, spacers 21 and 29 and covers 22 and 26 were laminated to achare to the base 2 having the above-described reaction layer 50 as shown in Figure 9, to produce the obscess sensor.

to form e reaction layer 50.

To the thus obtained glucose sensor, 10 µl of e mixed apueous solution of plucose and ascorbic add as sample liquid was supplied through sample supply ports 23 end 27. The supplied sample liquid immediately reached air ports 24 and 25 due to capillarity, and the reaction layer 80 on the main electrode system 19 was dissoluted.

A voltage of 4 1 V on the basis of the counter effects of 6 of the set decircles system? Now the mappined to the working decircle 8, and the current value I_g was the respect of the counter state of extent research. Since no codereductions and decircle counter of the counter

One minute after the supply of the sample (agait, a voltage of 4.6 ½ on the basis of the counter electhod's of the main electhode system 19 was applied to the working electhode 6 of the main electhode system 19, end the current value 1₃ after 5 excepts of the application was measured. The current value 1₃ was a total value of an outdistion current of potassium ferrographic openerated by a reaction to escorbic ocid and one genetated in a reduction of hubose by the GOD.

The emount of assorbic acid in the sample liquid was quantified from the current value I₀. A glucose concentration in the sample liquid was calculated from the quantified emount of ascorbic acid and the emount of potassium ferrooyanide obtained from the current value.

In this exemple, the main electrode system 19 and the sub electrode system 20 were provided on different sides from each other. Accordingly, the materials contained in the reaction layer 50 such as the GOD did not move onto the sub electrode system 20. As a result, there was no need to immobilize the GOD, potassium ferricyanide end the like to the main electrode system 19.

Apart from in the above-merisioned method, the base 2 can be produced by alemanting hes inelustring substrates 1 respectively bearing an electrode pattern on on surface thereof to each other. The structure of the sub-electrode system does not necessary correspord to that of the main electrode system. For examjet, the structure shown in Figure 10 can be used:

Example 6

Figure 11 is a sectional view of a glucose sensor produced as yet another example.

A base 2 was produced in the same menner as in Example 5.

A mixed aqueous solution of the GOD, potensium territopanide and the GMO was dropped on emain elsetrologues and the GMO was dropped on emain elsetrologues pattern 19 (e working electricide 6 and a counter electricide 7) of the base 2 and difficial to bern a rescribin layer \$1 in the same manner as in Example S. Next, o mixed aqueous solution of potession territogranide and the CMO was dropped onto a sub-electrode system 20 (a working electrode 8 and a counter electrode 9) and dried to form a reference layer 25 made from potassium ferritopanide and the CMO.

A spacer 3 and e cover 4 were integrally laminated on the base 2 in the same manner as in Example 5 to form the chaose sensor.

To the thus obtained plucose sensor, 10 µl of emixed aqueous solution of glucose and ascorbic acid as e-sample liquid was supplied through sample supply ports: 23 and 27. The sample liquid immediately reached air ports 24 and 28 due to carillitativ.

The reference layer 25 was dissolved in the sample ligited on the solvention system 20.0 Patiestims referreligated on the solvention system 20.0 Patiestims referreantide was reduced by assorbic acid in the sample layuid. A votage of 10.5 V on the basis of the counterelectricals 9 of the sub-electrical system 20 was applied to the working electrical 8 of the sub-electrical systems 20 ten accords after the supply of the sample ligati. A socurrent vision measured 5 seconds after the application was in proportion to the concentration of assorbic exist in the sample ligation.

The reaction layer 51 was dissolved in the sample injud on the main electrical system 19. Pottasium ferri-co-cyanida in the reaction layer 51 was changed into potassium ferrocyanida by two reactions: reduction by ascorbic ecid in the sample liquid and reduction in co-dicting glucose in the sample liquid by the GOD.

A voltage of +0.5 V on the basis of the counter electrode 7 of the main electrode system 19 was applied to the working electrode 6 of the main electrode system 19 one minute after the supply of the sample liquid to the sensor. The current value t₁ of 5 seconds after the application was measured.

The current value I₁ was a total value of the oxidetion current of potassium ferrocyanide generated by a reaction with ascorbic ecid end one generated by a reduction in oxidizing alucose by the GOD.

A concentration of ascorbic acid in the sample liquid was quantified from the response of the sub electrode system 20. A glucose concentration in the sample liquid was calculated from the quantified ascorbic acid concentration and the amount of potassium ferrocyanide obtained from the current value I.

An output current value in the sub electrical system 20 can not be accurately measured when the COD exists on the sub electrode system 20. The COD can be considerely prevented from moving onto the sub electrode system 20 by providing the main electrode system 18 and the sub electrode system 20 on the different surfaces of the substant is from each other as in the cample. No e result, a more accurate measurement can be entained.

Example 7

A saccharide sensor will be described as an example of the biosensor according to the present invention. Figure 12 is a sectional view of the seccharide sensor according to this example. A method for producing the se

A base 2 shown in Figure 12 was produced in the same manner as in Example 5.

Next, a mixed aqueous solution including the GOD as the oxidoreductises, potassium ferrifoyanide as the electron ecopotos and the CMC as the hydrophilic polymer was dropped on the main electrode system 19 and dried to form a first reaction layer 52. A mixed aqueous solution including the FDH as the

oxidoreductase, potassium ferricyandia es the electron ecogitors and the CMC as the hydrochilic polymer in e to phosphate buffer (pH = 4.5) was then dropped on the sub electrode system 20 and dried to form a second reaction layer 53. A spacer and e cover were laminated on the base 2 as in Example 5 to produce the saccharide sensor.

To the thus obtained seachwise sensor, 10 µ of placement and include and fluctose and fluctose are cample largely sens supplied photops in sample supply point 32 and 27. A vellage of the velocity of the 30 velocity of 10.00 velo

When the first end the second reaction layers 52 and 53 were dissolved in the sample liquid, the sub-

strates in the sample liquid were respectively coefficial with the oxideroclates operation to each layer. Potta-skim ferricyanide was reduced into potassium terrospanide was reduced into potassium terrospanide with an electron transiter in each layer. Next, by the application of the above predetermined voltages, an oxideonic current value corresponding to the generated action content value corresponding to the generated not applicable of the value of value of the value of value of the value of the value of va

The sensor response of the above seacharide sensor suptied with fruit juice as a sample liquid was measured. Glucosa and fructose in the fruit juice could be quantified.

The GOD used in the first reaction layer 52 and the FDH used in the second reaction layer 53 have different pH conditions for providing the highest enzyme reactivity from each other. Generally, the most appropriate pH condition often depends upon a kind of the used enzyme. When the first and the second reaction layers 52 and 53 are provided on the same surface of the substrata 1, a buffering component contained in the second reaction layer 53 moves into the first reaction layer 52 containing the GOD by the dispersion of the sample liguid. Thus, the most appropriate pH condition may not be obtained. Further, the enzyme may move onto a plurality of electrodes by the dispersion of the sample liquid. Tharefore, it is necessary to set a condition for not allowing the enzyme to move by immobilization or the like. As a result, the structure of the sensor can be limited

When the reaction leyers containing different encrymes are provided on the different surfaces of the insulating substants of time each other as in this example, a component in each reaction layer can be prevented from moving when each reaction layer is a disabled in the sample liquid. In this way, the pil on each electrode system can be easily extend to be the most appropriate to the entryme, and this encryme can be revel specified in the sample liquid.

In the above described examples, when the cover of and the spacer ere made from a transparent material such as a transparent synthetic resin, it is possible to observe the condition of the reaction layer end the introducing condition of the sample liquid in the passage from the outside.

In the foregoing examples, in order to supply the sample liquid to the reaction layer more smoothly, a lecithin layer may be formed by developing an organic solvent solution of lecition through the sample supply port into the reaction layer and drying thereof.

When the lecithin layer is provided, the sample liquid can be supplied even when the passage defined by the base, the cover and the spacer is not small enough to cause confliatity.

Since the amount of the sample liquid to be supplied depends upon the capacity of this passage, there is no need to previously quantity it. In addition, the evaporation of the sample liquid during the measurement can be minimized, thereby attaining a more accurate measurement.

The sample supply port is not necessarily distinguishable from the air port. It is possible to supply the

sample liquid through the eir port, using the sample supply port as an air port.

The oxideworks are the GOD in the coop.

The outdoreductions such as the GOD in the resontion layer in not expectedly immobilized to he main electrode system. However, since the residen layer contains the hydrochile polymer, the eligenmic of the metherial is prevented due to increased viscosity of the anaptie figuid when the reaction layer is discorted in the sample faguid. Therefore, the material mixing up the restorict layer does not move cost the able electrical form of the production of the reciproc can be effectively introduction of the reciproc can be effectively mixing the conduction among reliable measurement.

When he sample logical is supplied through his sample supply only. It is effective to provide the sub-electrode system in the vicinity of the sample supply once to their sample supply proceeds bowed the size post through the sample supply proxi Therefore, a proaction toward the sub-electrode system that is promoted to the sub-electrode system that is promoted to the sub-electrode system that is promoted toward the sub-electrode system that is promoted toward to the sub-electrode system, this does not cause any problem.

In Examples 5, 6 and 7, the sub electrode system has the same electrode pattern as the main electroda system. However, it does not have to be the same. For example, the pattern shown in Figure 10 can be used.

There is no need to form all of the reaction layers, the first reaction layer, the second reaction layers, the first reaction layer at the telectrode systems as in the foregoing accumpter. When the base, the spacer and the cover ere integrated, the reaction layer may be formed on a reverse surface of the cover and the like so as to face the nesseon.

Further, in the above-described examples, a method for quantifying glucose and fructose is shown. However, the present invention can be widely used in systems using an enzyme reaction, as en alcohol sensor, a factic acid sensor, a cholesterol sensor and an amino acid sensor.

Moreover, in the foregoing exemples, the GOD and the FDH ara used as the oxidoreductase. However, alcohol oxidase, lactase oxidase, lactase oxidase rate, cholesterol oxidase, xanthine oxidase and emino acid oxidase and the like can be used as well.

The hydrophic opymer is not limited to the CMC used in the examples. Other collabora derivatives such as hydroxy ethyl celetiose, hydroxy propyl celetore, methyl cellulose, shydroxy propyl celetore, methyl cellulose, shydroxy ethyl celetore can be used. Norcover, the same effect can be statiened by using polyminghymolicane, polyming alloohol, getalin or its derivative, and of the sales. The method of the sales.

salts, starch or its derivatives and maleic anhydride or its salts.

its salts.

As electron ecceptors, apart from potassium ferricytanide used in the above-mentioned examples, p-benzoguingne, phenezinemethosulfate, methylene blue s

and ferrocene derivatives can be used.

Moreover, in the foregoing examples, the coldoreductase end the electron acceptors are dissolved in the semple liquid. However, they may be immobilized to be

insoluble in the sample liquid.

The above-described efectrode system is not limited to e two-electrode system having only a working electrode and a counter electrode. A three-electrode system, including an additional reference electrode may be used, so that more procise values ere obtains-

Claims

1. A biosensor comprising:

en electrical insulating substrate (1) having a plurality of surfaces.

a plannily of electrode systems (19, 20) formed on a at least two of the plurality of surfaces of the substrate (1), each of the electrode systems (19, 20) having a working electrode (6, 8)

and a counter electrodo (7, 9), and a plurality of reaction layers (52, 53) provided on at least two of the plurality of surfaces of the substrate (1), each of the reaction layers (52, 53) containing an oxidoreductase.

wherein the plurality of electrode systems (19, 20) are respectively provided on different surfaces of the substrate (1), the reaction leyers as (25, 55) are provided on the surfaces on which he plurality of electrode systems (19, 20) are formed, end the kinds of oxidoreductuses contained in the plurality of reaction layers (52, 55).

 A blosensor according to dalm 1, wherein the substrate (1) is plate-shaped and wherein one (19) of the plurality of electrode systems is provided on one surface of the substrate (1) and another electrode system (20) is provided on another surface of the substrate (1).

are different from each other

- A biosensor eccording to claim 1 or 2, wherein the oxidoreductase is selected from the group consistor of functione dehydrogenase, glacose acidase, elcohol oxidase, lactase oxidase, lactase dehydrogenase, chalesterol oxidase, xarritine oxidase end amino acid oxidase.
- A biosensor eccording to any of the preceding claims, wherein each of the pursity of reaction layers (52, 53) further contains an electron ecceptor and a hydrophilic polymer.

- 5. A biosensor occording to claim 4, wherein the hydrophilic polymer is selected from the group consisting of carbony methyl cellulose, hydroxy peopl cellulose, hydroxy peopl cellulose, methyl cellulose, ethyl cellulose, polymyn derivalose, polymyn derivalose, gelation or its deviraleys, anytic acid or its salts, methacnylia caid or its salts, sterch or its deviraleys, and make anythicid or its salts.
- A biosensor eccording to claims 4 or 5, wherein the electron acceptor is selected from the group consisting of potassium ferricyanide, p-benzoquinone, phenazinemetho sulfate, methylene blue and ferrocene.
- A method for quantifying e substrate contained in e sample liquid wherein e biosensor according to any of the precoding claims is used to quantify at least two different substrates.

Fig. 1

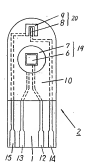
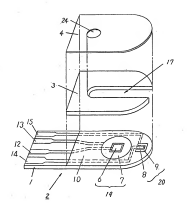
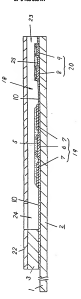


Fig. 2





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Fig. 4

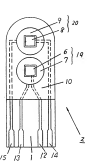


Fig. 5

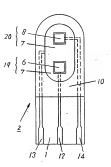


Fig. 6

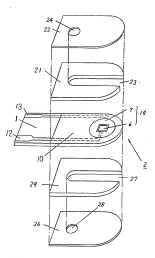


Fig. 7

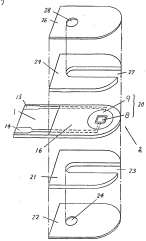


Fig. 8

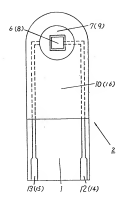


Fig. 10

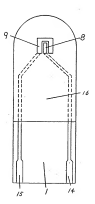


Fig. 11

Fig. 12

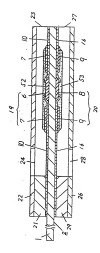


Fig. 13

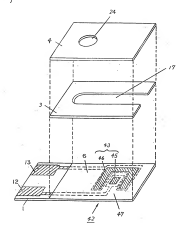
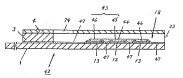


Fig. 14



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European Palent EUROPEAN SEARCH REPORT EP 96 10 8449

Category	Citation of document with i	adication, where appropriats,	Reterrent to chies	CLASSIFICATION OF THE APPLICATION (BLCLS)	
A	EP-A-0 127 958 (GEN INC.)	ETICS INTERNATIONAL,	to chan	G01N27/327 C12Q1/00	
D,A	EP-A-0 359 831 (MAT INDUSTRIAL CO.)	SUSHITA ELECTRIC		C12Q1/26	
),P,	EP-A-0 582 504 (MAT INDUSTRIAL CO; LTD.	SUSHITA ELECTRIC			
				TECHNICAL PRELOS SECARCIED (SECLE)	
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	The present search report has b	con drawn up for all claims Date of completion of the search	_		
THE HAGUE		17 July 1996	Car	tagena y Abella,P	
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